Europäisches Patentamt

European Patent Office

Office européen des brevets



(11) EP 1 063 244 A2

(12)

EUROPEAN PATENT APPLICATION

- (43) Date of publication: 27.12.2000 Bulletin 2000/52
- (21) Application number: 00121154.9
- (22) Date of filing: 18.12.1996
- (84) Designated Contracting States: BE DE ES FR GB IT NL
- (30) Priority: 19.12.1995 US 8893 P
- (62) Document number(s) of the earlier application(s) in accordance with Art. 76 EPC: 96944998.2 / 0 868 445
- (71) Applicant: EXXON CHEMICAL PATENTS INC. Baytown, TX 77520-2149 (US)
- (72) Inventors:
 - Turner, Howard W.
 Campbell, CA 95008 (US)
 - Speed, Charles S. Dayton, Texas 77535 (US)
 - Folie, Bernard J.
 1380 Lasne (BE)

- (51) Int Cl.⁷: **C08F 210/18**, C08F 2/06, C08F 4/645
 - Crowther, Donna J.
 Seabrook, Texas 77586 (US)
 - Walzer, John F.
 Seabrook, Texas 77586 (US)
 - Fisher, Richard A.
 Malvern, Pennsylvania 19355 (US)
 - Vaughan, George A. Houston, Texas 77025 (US)

1830 Machelen (BE)

(74) Representative: Veldhuizen, Albert Dirk Willem et al Exxon Chemical Europe Inc., P.O.Box 105

Remarks:

This application was filed on 29 - 09 - 2000 as a divisional application to the application mentioned under INID code 62.

(54) High temperature olefin polymerization process

(57) The invention provides a method for insertion polymerization of olefinically or acetylenically unsaturated monomers comprising contacting one or more of said monomers under suitable polymerization process con-

ditions with an ionic catalyst composition comprising the reaction product of a catalytically suitable Group 4, 5, 6 or 8 transition metal compound having a univalent hydride, alkyl or dilyl ligand and a hydrated salt comprising a Group 1 or 2 cation and a non-coordinating anion.

Description

5

10

25

30

50

Field of the Invention

[0001] This invention relates to catalyst system selection and high temperature process conditions for polymerization of monomers containing ethylenic or acetylenic unsaturation wherein the catalyst system is single-sited and comprises a transition metal cation and a stabilizing, compatible non-coordinating anion.

Background of the Invention

[0002] Ionic catalyst systems for olefin polymerization based on transition metal compounds such as metallocene compounds are recent but now well-known. Such metallocene compounds are based on transition metals capable of supporting at least one pi-bound aromatic ligand, typically a cyclopentadienyl or substituted cyclopentadienyl ligand and at least one additional ligand which can be abstracted so as to form a catalytically active cation structure. Cocatalyst anions are needed to stabilize these cations, which are highly reactive, but those anions must be capable of doing so without either closely coordinating with the cation that its reactivity, and thus polymerization activity, is diminished, or transferring an anionic fragment to the cation so as to alter its polymerization capability. Single coordination complexes and multi-nuclear coordination complexes based on the Group IIIA or 13 elements, such as boron or aluminum, have shown particular suitability as effective co-catalyst anion sources. See, the disclosures in EP-A-0 277 003, EP-A-0 277 004 and their equivalents, U.S. patents 5,198,401 and 5,278,119, particularly Examples 27 and 32. Example 32 of '119 discloses the use of bis(trimethylsilylcyclopentadienyl) hafnium dimethyl as the metallocene compound. Example 27 of '401 lists dimethylsilylbis (indenyl) zirconium dimethyl in column 9, line 16.

[0003] Homogeneous processes such as high pressure, high temperature polymerization processes, such as those at 500 bar and above, have shown particular suitability with such catalyst systems since these systems exhibit greater stability than those using alumoxanes co-catalysts at 160 °C and above. This allows for the greater productivity that occurs with greater reactivities at high temperature, the high pressure maintaining an essentially single phase reaction medium while permitting the higher temperatures. See U.S. patent 5,408,017 and equivalent WO-A-93/05732. A generic, broad description of suitable metallocene compounds is given as is a generic description of suitable anion precursors. The examples include the metallocene compounds dimethylsilylbis (4,5,6,7 tetrahydro-indenyl) zirconium dimethyl and dimethylsilylbis (4,5,6,7 tetrahydro-indenyl) hafnium dimethyl. The former showed significantly higher catalyst productivity (expressed as kg-PE/g-activator) in Table 2, ranging from 100 to 160, while that of the hafnium compound exhibited only 60. A similar olefin polymerization process that can be operated at high temperature and pressure is disclosed in WO 95/07941. The principal problem addressed is build up of polar material in recycle streams of the reactor, the solution is the use of bulky scavengers, e.g., those having at least one tertiary carbon atom, such as triisobutyl alumoxane. Me₂Si(Ind)₂Hf Me₂ is shown in Comparative Example 3 and Example 4, and said to be active longer and illustrative of high conversions.

[0004] The selection of substituents on the *pi*-bound, cyclopentadienyl ligands in the metallocene compounds has been identified as a means of increasing performance in olefin polymerization processes. For example, see U.S. patent 5,304,614 where specifically substituted indenyl ligands and their methods of preparation are described. Very high molecular weight polyethylene at high activity is said to be possible by use of the described metallocene compounds based on any of the group IVB, Vb or Vlb metals. Zirconium and hafnium are said to be preferred and the preferred substituent structure is characterized by 1) alkylene or silylene groups bridging two cyclopentadienyl ligands bound to the metal atoms and 2) 4,7-substituted benzo groups fused on the respective sides opposite the H- or C₂ to C₄ 2-substituent on those same cyclopentadienyl groups. Examples 14-18 are of homopolymer polyethylene using specifically, substituted bis(indenyl)zirconocenes with an alumoxane co-catalyst at a temperature of 70 °C and ethylene pressure of 5 bar. Productivity ranged from 35 to 56 g PE for 1 hour of polymerization reaction. The highest production was reported for Me₂ Si (3,4,7-Me₃lnd)₃ Zr Cl₂.

[0005] In EP-A1-0 612 768, bridged and unbridged hafnium metallocene compounds stabilized by non-coordinating anions after activated with alkyl-aluminum co-catalysts are said to demonstrate high catalyst activity over the zirconium analogues when utilized in processes at temperatures at or exceeding 120 °C. The medium pressure solution processes are preferably to be conducted at pressures of 500 to 3500 kg/cm². All listed hafnocenes are dichloride-substituted embodiments and each of the working examples of the invention is alkylene bridged, with cyclopentadienyl, indenyl or fluorenyl pi-bound ligands.

[0006] Catalyst systems based on monocyclopentadienyl titanium compounds activated with alumoxane suitable for the preparation of ethylene-α-olefin copolymers of high molecular weight and high a-olefin content are described in U. S. patent 5,264,405. This patent teaches that the cyclopentadienyl group should be fully substituted with methyl groups and bridged to an amido group having an aliphatic or alicyclic hydrocarbyl ligand bonded through a 1° or 2° carbon. Copolymerization of ethylene with propylene in Example 45 with a bridged monocyclopentadienyl Group 4 metal cat-

alyst compound at 80 °C produced a copolymer with 20 wt.% ethylene having an M_n of about 20,080. In each Example 55 with the same catalyst as with Example 45, at a reaction temperature of 140 °C, an ethylene-propylene copolymer having a density of 0.863, indicative of an amorphous ethylene copolymer, exhibited an M_n of about 46,500.

[0007] Due to the sensitivities of ionic catalyst systems to polar impurities, solution polymerization processes utilizing scavenging compounds, for example alkyl aluminum compounds or alkyl alumoxanes, have been described. See, for example, U.S. patents 5,153,157 and 5,241,025, describing Group-IIIA metal scavenger compounds and processes. WO-A-94/07927 addresses a similar process as adapted for monocyclopentadienyl catalyst systems, it describes advantages of bulky scavengers when the monocyclopentadienyl catalyst compounds do not contain bulky substituents to impede interaction with the scavenging compounds. Triisoprenyl aluminum is exemplified and triisobutyl aluminum is listed along with tri-n-octyl and tri-n-hexyl aluminum as suitable bulky scavengers.

[0008] In view of the breadth of the disclosures concerning the use of metallocenes for olefin polymerization, and the general industrial need to employ the most effective catalysts and processes so as to have increased productivities while maintaining both comonomer incorporation and high molecular weight polymer preparation capability, additional investigative efforts were required. In particular general knowledge based on observations in the field gave rise to the traditional understanding that under temperatures exceeding about 80 °C increased comonomer incorporation using metallocene catalysts resulted in lower molecular weight polymer.

Invention Disclosure

10

15

20

30

35

40

[0009] It has been discovered in resulting investigations that even when subjected to the demanding conditions of high temperature in solution and high pressure processes, high molecular weight ethylene copolymers (M.I.≤ 10, preferably, M.I. ≤ 1) can be polymerized in the presence of stable, single-sited discrete ionic catalyst systems, for example, those having cations derived from the transition metal compound group consisting of 1) bridged, unsubstituted- or substituted-indenyl, or fluorenyl group containing, hafnium compounds; 2) bridged hetero-atom containing, substituted or unsubstituted monocyclopentadienyl titanium compounds; 3) unbridged bulky Group 15 containing, bulky monocyclopentadienyl titanium compounds; 4) Group 4 or 5 compounds containing bulky chelating diamide ancillary ligands; and 5) Group 8 metal diimine compounds wherein the metal is in a -2 oxidation state, such as Pd(II) and Ni(II). Accordingly the invention is in part a process for copolymerizing ethylene and at least one higher olefin, that is C₃ to C₂₀, preferably C₄ to C₈, alpha-olefin, diolefin, or C₄ to C₂₀ cyclic olefin, comprising the step of contacting said monomers with one or more of the stable, single-sited ionic catalysts with a cation above and a stabilizing, compatible non-coordinating anion in a polymerization reaction with a reaction medium temperature of 90 °C or above, and optionally at a pressure exceeding 50 bar, preferably exceeding 75 bar for solution processes and 500 bar for high pressure supercritical phase processes. The invention is also in part a modified process comprising the additional step of introducing into either the reaction medium, recycle stream or the monomer feedstocks prior to the polymerization reaction, a scavenger for polar impurities, most preferably a trialkyl aluminum each alkyl being a long-chain, linear-alkyl.

Detailed Description and Best Mode of the Invention

[0010] The bridged hafnium compounds of the invention include those having one or more carbon, silicon, or germanium atoms bridging two cyclopentadienyl (Cp) ligands of the hafnium metal centers, said ligands optionally containing one or more hydrocarbon substituents. When the Cp ligand is an indenyl or fluorenyl group, substitutions can be made either on the 5 or 6 member ring carbon atoms. Substituents typically include one or more C₁ to C₃₀ hydrocarbon groups selected from linear, branched, cyclic, aliphatic, aromatic or combined groups, whether in a fused-ring or pendant configuration. For the purposes of this application the term "hydrocarbon" is meant to include those compounds or groups that have essentially hydrocarbon characteristics but optionally contain not more than about 10 mol. % non-carbon, polar atoms, such as oxygen, sulfur, nitrogen and phosphorous. Similarly the use of hetero-atom containing cyclopentadienyl rings, where a non-carbon atom replaces one of the ring carbons, is considered for this specification to be within the terms "cyclopentadienyl", "indenyl", and "fluorenyl".

[0011] Specific bridged hafnium catalysts include those derived from: (1) indenyl-based complexes such as the *rac*-or *meso*- isomer of dimethylsilyl bis (indenyl)hafnium dimethyl, dimethylsilyl bis(4,5,6,7- tetrahydro-indenyl) hafnium dimethyl, dimethylsilyl bis(2-propyl-indenyl) hafnium dimethyl, dimethylsilyl bis(4-methyl, 2-phenyl-indenyl) hafnium dimethyl; (2) cyclopentadienyl complexes such as dimethylsilyl (cyclopentadienyl)(tetramethyl cyclopentadienyl) hafnium dibenzyl, dimethylsilyl bis(cyclopentadienyl) hafnium dimethyl; and (3) fluorenyl-based complexes such as dibutylsilyl (fluorenyl) (cyclopentadienyl) hafnium dimethyl and dimethylsilyl (indenyl) (fluorenyl) hafnium dimethyl.

[0012] In particular, for the bridged bis indenyl hafnium compounds, it has been found that increasing the degree of substitution on the indenyl ligands is effective for increased comonomer incorporation, an effect surprising in view of the general knowledge in the art. Thus when the indenyl substituents include at least two sigma bound hydrocarbon

radicals replacing hydrogen atoms on the ring atoms, the performance exceeds that where either of zero substituents or just one substituent are present. For example dimethylsilyl bis(2-methyl, 4-phenyl-indenyl) hafnium dimethyl has been found to provide one or both of higher comonomer incorporation and higher molecular weight as compared to dimethylsilyl bis(2-methyl-indenyl) hafnium dimethyl, which is better in one or both of these features than dimethylsilyl bis(indenyl) hafnium dimethyl. Thus preferably the ligation on the indenyl radicals in the bulky hafnium compounds will generally comprise two or more C₁ to C₃₀ hydrocarbon substituents as defined above.

[0013] The silicon bridged hetero-atom containing, substituted or unsubstituted monocyclopentadienyl titanium compounds of the invention are any of those described in the art, see for example those described in U.S. patent 5,264,405, WO 92/00333 and U.S. patent 5,408,017. The substituted cyclopentadienyl ligand is one having hydrocarbon substituents as defined for the cyclopentadienyl, indenyl and fluorenyl ligands of the bridged hafnium compounds above. Each document in incorporated by reference for purposes of U.S. patent practice.

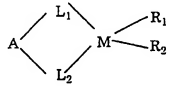
[0014] The unbridged, bulky Group 15 containing, bulky monocyclopentadienyl titanium compounds of the invention are unbridged titanium compounds having ancillary ligands including a substituted, bulky cyclopentadienyl ligand, a substituted, bulky Group 15 heteroatom ligand, and two uninegative, activation reactive ligands at least one of which that can be abstracted for activation of the remaining metal compound to a catalytically active state and one of which is either similarly abstractable or has a σ-bond to the transition metal into which an olefin or diolefin can insert for coordination polymerization. Typically, for any of the catalysts of this invention, uninegative, activation reactive ligands are selected from the group comprising hydride, lower alkyl, e.g., C₁ to C₄, or silyl. The bulky cyclopentadienyl ligand is one having substituents as defined for the cyclopentadienyl, indenyl or fluorenyl ligands of the bridged hafnium compounds above, particularly lower alkyl-substituted ligands, such as tetramethyl and pentamethyl cyclopentadienyl ligands. The bulky Group 15 heteroatom ligand is typically a hydrocarbon substituted Group 15 element, preferably nitrogen, wherein the hydrocarbon substituents are covalently bound to the heteroatom through a secondary or tertiary carbon or silicon atom. See also, copending application U.S. Serial Number 08/545,973, filed October 20, 1995 (Atty. Docket No. 958059) which is incorporated by reference for purposes of U.S. patent practice.

[0015] Ionic catalysts derived from Group IVB metal precursors where the ancillary ligand system does not contain a cyclopentadienyl ligand may also be employed in this invention. In general, these systems will be prepared from an ionic activator or Lewis Acid activator and a Group IVB metal containing precursor of the following general formula:

30

35

25



wherein:

40

A is an optional bridging substituent,

L₁ and L₂ are the same or different non-Cp ancillary ligands,

M is a Group IVB metal, and

R₁ and R₂ are the same or different σ-bonded groups such as hydride or hydrocarbyl.

45

50

55

[0016] An example of a suitable non-Cp ancillary ligand system is described in "Conformationally Rigid Diamide Complexes: Synthesis and Structure of Tantalum(III) Alkyne Derivatives", D. H. McConville, *et al, Organometallics* 1995, *14*, 3154-3156. Among the Group 4 metals, the Group 4 metals are preferred especially those of titanium. The Group 4 metal compounds will additionally comprise at least two uninegative, activation reactive ligands at least one of which that can be abstracted for activation of the remaining metal compound to a catalytically active state and one of which is either similarly abstractable or has a σ-bond to the transition metal into which an olefin or diolefin can insert for coordination polymerization. The Group 4 metal compounds having the described ligands can be prepared as illustrated in the *Organometallics* article except that the Group 5 metal halide is replaced with a Group 4 tetrahalide [0017] The Ni²+ and Pd²+ complexes of the invention are diimine complexes that can be prepared by methods equivalent to those used for the compounds described in "New Pd(II)- and Ni(II)- Based Catalysts for Polymerization of Ethylene and α-Olefins", M. Brookhart, *et al, J. Am. Chem. Soc.*, 1995, *117*, 6414-6415. These can be either the dialkyl ether adducts as described, or alkylated reaction products of the described dihalide complexes that can be activated to a cationic state by the anion precursors according to the invention.

[0018] In general effective scavenging compounds for ionic polymerization catalysts include those aluminum alkyl compounds listed in U.S. patents 5,153,157 and 5,241,025. The term "scavenger" is used in its art-recognized sense of being sufficiently Lewis acidic to coordinate with polar contaminates and impurities adventiously occurring in the polymerization feedstreams or reaction medium In particular, for processes utilizing recycle streams of unconverted monomer for reprocessing, the necessity to use polar compounds as catalyst deactivators, or "killers", such as water or lower alcohols, effectively necessitates the use of scavengers, as does the natural occurrence of polar impurities in monomer feedstreams. It has been discovered, however, that the use of excess scavenger has a deleterious effect on catalyst performance, and that the proper choice of scavenger is important to minimize the effects Addition of excess scavenger causes lower productivity, molecular weight and comonomer incorporation. This effect can be easily seen by comparing Examples 1.2, 1.6, 1.7, and 1.8 in Table 1 of the examples section, when the addition of 6 equivalents of triethylaluminum (TEAL) caused an increase in productivity but as more TEAL was added productivity decreased as did the molecular weight and comonomer content of the product. Replacing TEAL with triisobutyl aluminum (TIBA) (Examples 1.9 - 1.12) improved the catalyst performance under excess scavenger conditions but did not completely remedy the situation.

10

15

20

30

35

40

55

[0019] Experiments under high temperature conditions showed similar trends. In order to further improve the performance of the ionic catalyst system, a systematic study of more substituted scavengers was carried out. Based on the observation that TIBA performed better than TEAL, it was expected that more highly substituted aluminum alkyl compounds, such as Al(CH₂CMe₃)₂Me, where Me is methyl, would be superior to TIBA because of the increase steric bulk around the Al atom. Comparative tests under high temperature conditions showed that Al(CH₂CMe₃)₂Me was inferior to TIBA and caused sudden loss of productivity at very low levels of scavenger. Further studies showed that the preferred scavenger is a long chain, linear tri-alkyl aluminum compound, and that longer chains are preferred over shorter chains.

[0020] Non-limiting examples of effective long chain, linear tri-alkyl ligand-containing scavengers include those comprised in the group defined by the formula M'R'R", where M' is AI, and each of the R groups independently is a C_4 or higher linear, branched or cyclic alkyl group, preferably C_6 or higher, most preferably C_8 or higher. The long chain, linear alkyl aluminums where each alkyl substituent was of a length of C_8 or higher, preferably C_9 and higher were observed to exhibit optimal performance, that defined as having the least deleterious effect when used at a level in excess of the optimum level as described in the following paragraph. Specifically included are: tri-n-octyl aluminum, tri-n-decyl aluminum, tri-n-decyl aluminum, tri-n-hexadecyl aluminum, and the higher carbon number equivalents, e.g., $(C_{20})_3$ AI, including those with mixed ligation, and mixed scavenger compounds as well. The hydrolyzed derivatives of these alkyl-ligand containing oreanoaluminum compounds will additionally be suitable Additionally, it will be apparent that those scavenging compounds comprising both long-chain, linear and bulky ligands or mixed linear ligands, each ligand as described above, will also be suitable, but perhaps less desirable due to more involved or expensive syntheses.

[0021] The long chain, linear-alkyl ligand-containing scavengers of the foregoing paragraph will be useful in any insertion polymerization process for olefinically or acetylenically unsaturated monomers with any of the ionic catalyst systems known in the art or those in development, where such make use of non-coordinating anions and the resulting catalyst sensitivities require elimination of polar impurities. Suitable catalyst systems appear in the catalyst references listed in the Background in this application, additional catalysts include those Group 5 and 6 transition metal catalyst systems of WO 94/01471 based on U.S. Serial Number 08/086,772 filed July 1, 1993, the disclosures of which are incorporated by reference for purposes of U.S. patent practice. Examples of suitable process conditibns include those of gas phase, solution, slurry or bulk polymerization processes for any polymers or copolymers of two or monomers selected from the group consisting of ethylene, propylene, C₄-C₂₀ α-olefins, strained ring cyclic olefins, macromers of up to 100 or more mer units having olefinic unsaturation in the 1-position, or acetylenically unsaturated monomers. Such processes utilize -50 °C to 300 °C temperature and 0 to 3000 bar pressure. Polymers and copolymers having molecular weights equivalent to an M.I. of 100 and below can be prepared in these processes.

[0022] The scavengers, whichever are selected, should be utilized in a manner consistent with the sought productivity, polymer molecular weight, and polymer comonomer content. In particular, only that amount sufficient to neutralize the effects of the adventitious catalyst poisons should be utilized, the better purified the comonomer feedstreams into the reactor and as well the other feedstreams or added reactants and recycle, the less scavenger will be required. The amount is preferably as little as possible and can be adjusted empirically by observing the rates of reaction, adiabatic temperature rise, and other indicators of reaction efficiencies observable in the polymerization process, each being maximized holding all variables other than the addition of scavenger constant. Excess scavenger has been observed to cause a reduction in productivity, molecular weight and comonomer incorporation.

[0023] Means of preparing the ionic catalyst systems comprising cations of the described transition metal compounds and suitable non-coordinating anions are conventionally known, see for example U.S. patent 5,198,401 and WO 92/00333. Typically the methods comprise obtaining from commercial sources or synthesizing the selected transition metal compounds comprising an abstractable ligand, e.g., hydride, alkyl or silyl group, and contacting them with an

ionizing non-coordinating anion source or precursor in an aromatic solvent. The anion source acts to ionize the transition metal compounds by abstracting the univalent hydride, alkyl or silyl ligand that complements its total valency. The abstraction leaves the transition metal compounds in a +1 cationic state, which is counterbalanced by the stable, compatible and bulky, non-coordinating anion. See the fuller description in U.S. patent 5,198,401, referred to above. Conditions of adiabatic polymerization processes such as practiced at high pressure, result in temperatures exceeding 160 °C and above. These higher operating temperatures lead to increasing instability that is reflected in more difficult higher olefin comonomer incorporation and reduced molecular weight polymer. However, the single site catalysts of this invention under those same conditions evidence higher stability, greater comonomer incorporation and higher retained molecular weight as evidenced in the tabulated observations below.

10

20

25

30

35

55

[0024] The non-coordinating anion sources of the invention includes any of those conventionally known to be useful for olefin polymerization with metallocenes, including those known to be useful with the single cyclopentadienyl containing Group 4 transition metals. A representative listing of suitable anions is in U.S. 5,198,401, EP-A-0 426 637, EP-A-0 427 697, EP-A-0 520 732, EP-A-0 573 403, WO 95/24268 and U.S. patent 5,387,568. Those may be introduced into the catalyst preparation step as either ionic compounds having a cation which abstracts a non-cyclopentadienyl ligand of the transition metal compounds or as neutral compounds which upon abstraction of the non-cyclopentadienyl ligand, leave as a by-product the non-coordinating anion portion. Additionally, it is known that the use of alkylating compounds along with an anion source enables the use of transition metal compounds having ligands too strongly bound to the transition metal center to be abstracted by the anion source, e.g., transition metal dihalides. Typical alkylating sources may be any of the strongly Lewis acidic organoaluminum compounds such as the lower carbon number alkyl aluminums and alkylalumoxanes. See EP-A-0 500 944, EP-A-0 570 982 and EP-A1-0 612 768 for in situ processes describing the reaction of alkyl aluminum compounds with dihalo-substituted metallocene compounds prior to or with the addition of activating anion precursor compounds. Each of the above documents is incorporated by reference for purposes of U.S. patent practice. Most preferably transition metal compounds not having halide ligands on the metal center are used for the ionic catalyst systems of this invention since in situ alkylation processes may result in competing reactions and interactions that tend to interfere with the overall polymerization efficiency under conditions of high temperature in accordance with this invention.

[0025] A class of preferred anion precursor compounds are hydrated salts comprising a Group 1 or 2 cation and a non-coordinating anion as described above. The hydrated salts can be prepared by reaction of the metal cation-noncoordinating anion salt with water, for example, by hydrolysis of the commercially available or readily synthesized LiB (pfp)₄ which yields [Li • H₂O] [B(pfp)₄], where (pfp) is penta- or perfluorophenyl. The by-product of ionization of the abstractable ligand-containing transition metal compounds with this precursor is LiOH which is non-volatile and thus is not recycled in the separation phase wherein the polymer is removed from monomer and any diluent prior to recycle. Tests have shown that this low-cost precursor operates to form ionic catalysts having properties essentially equivalent to those formed with the preferred precursor compounds represented by [Ph3C][B(pfp)4] and [PhMe2NH][B(pfp)4], Ph representing phenyl and Me representing methyl. As with the long chain, linear-alkyl ligand-containing scavengers noted above, this non-volatile by-product anion precursor will be suitable for use with any of the ionic catalyst systems known in the art or those in development, where such make use of non-coordinating anions. Catalytically suitable transition metal compounds capable of cationization include those Group 4 to 6 and 8 compounds addressed above. Again, examples of such include gas phase, solution, slurry and bulk polymerization processes for any polymers or copolymers of two or monomers selected from the group consisting of ethylene, propylene, C₄-C₂₀ α-olefins, strained ring cyclic olefins, macromers of up to 100 or more mer units having olefinic unsaturation in the 1-position, or acetylenically unsaturated monomers. Such processes utilize -50 °C to 300 °C temperature and 0 to 3000 bar pressure.

[0026] Known alkylalumoxanes are additionally suitable as catalyst activators for the invention single-site transition metal compounds comprising halide ligands, however these are less preferred since the alumoxane activators are less temperature stable at or above about 160 °C. The alumoxane component includes all those useful as a catalyst activator, typically such will be an oligomeric aluminum compound represented by the general formula (R-Al-O)_n, which is a cyclic compound, or R(R-Al-O)_nAlR₂, which is a linear compound. In the general alumoxane formula R is a C₁ to C₅ alkyl radical, for example, methyl, ethyl, propyl, butyl or pentyl and "n" is an integer from 1 to about 50. Most preferably, R is methyl and "n" is at least 4. Alumoxanes can be prepared by various procedures known in the art. For example, an aluminum alkyl may be treated with water dissolved in an inert organic solvent, or it may be contacted with a hydrated salt, such as hydrated copper sulfate suspended in an inert organic solvent, to yield an alumoxane. Generally, however prepared, the reaction of an aluminum alkyl with a limited amount of water yields a mixture of the linear and cyclic species of the alumoxane.

[0027] A preferred process -of polymerization is that conducted at high pressure, that is at from 200 to 3000 bar, preferably from 500 to 2500 bar in a homogeneous single phase or two fluid phases, with or without unreactive diluents or sulvents at temperatures generally above the melting point of the polymer being produced. Such processes are typically known and may include the use of scavenger compounds and catalyst deactivation or killing steps, see for example U.S. patent 5.408.017, WO 95/07941, and WO 92/14766. Each of these documents and their U.S. counter-

parts are incorporated by reference for purposes of U.S. patent practice. Preferred catalyst deactivators, or killers, include high molecular weight, non-recyclable compounds, such as poly vinyl alcohol which exhibit the functional capacity to complex with the catalysts so as to deactivate them while not forming volatile polar by-products or residual unreacted compounds.

[0028] Another preferred process in which any of the catalyst, cocatalyst and scavenger selections disclosed in this application can be advantageously practiced is that of a continuous, solution process operated at or above 90 °C to 120 °C, even above 150 °C or above 160 °C, up to about 300 °C. Typically this process is conducted in an inert hydrocarbon solvent, linear, cyclic or branched aliphatic, or aromatic, at a pressure of from 20 to 200 bar. The disclosures of U.S. patent applications Ser. No. 08/426,363, filed April 21, 1995, and Serial Number. 08/545,973 (Atty. Docket 95B059), filed October 20, 1995, provide relevant description, even when operated at the elevated temperature ranges under supercritical conditions disclosed in this application. These documents also are incorporated by reference for purposes of U.S. patent practice.

[0029] For optimal polymerization results the processes should be designed or conducted such that the cocatalyst components, that is the transition metal compounds and the anion precursor compounds, are maintained separately until just prior to or during polymerization use in the chosen reactor. An example is the use of dual injection of each catalyst component directly into the reactor or the use of T- or multi-joint mixing chambers just prior to injection into the reactor. Alternatively the catalyst may be formed *in-situ* by the independent addition of ionic activator, ligand stabilized metal halide, and scavenger directly into the reactor or the use of T- or multi-joint mixing chambers just prior to injection into the reactor. Additional optimization can be achieved when the scavenger compound is introduced into the reactor independently of the catalyst system, or compounds.

[0030] Though directed specifically for high temperature, particularly high pressure or solution, processes, it will be apparent that the use of heterogeneous support material such as polymeric and metal or metalloid oxide supports will enable slurry or gas phase use of the process steps and components described and will likely achieve similar benefits of higher productivities, higher molecular weights, and higher levels of comonomer incorporation for the ethylene copolymer products capable of production using the disclosed copolymerizable monomers. Illustrative support methods appear in WO 91/09882, WO 94/00500, WO 94/03506, WO 94/07928, and in WO 96/04319 based on U.S. application Serial Number 08/474,948 filed June 7, 1995, in turn based on U.S. Serial Number 08/285,380 filed August 3, 1994. A suitable slurry process is described in U.S. patent 5,229,478. Each of the foregoing are incorporated by reference for purposes of U.S. patent practice.

[0031] The following examples are presented to illustrate the foregoing discussion. All parts, proportions and percentages are by weight unless otherwise indicated. Although the examples may be directed to certain embodiments of the present invention, they are not to be viewed as limiting the invention in any specific respect. Methods of determining MW and monomer contents by GPC and NMR for the illustrative EPC examples of the invention are described in U.S. patent 5,229,478 which is incorporated by reference for purposes of U.S. patent practice. For the following examples certain abbreviations have been utilized for convenience: Cp (cyclopentadienyl), Me(methyl), Ind (indenyl), Ph (phenyl), pfp (pentafluorophenyl), r-(racemic), Et (ethyl), Cp* (permethylated cyclopentadienyl), Cod (cyclododecyl), TOA (tri-n-octyl aluminum), TEAL (triethyl aluminum) and TIBA (triisobutyl aluminum).

EXAMPLES

20

30

40

50

Part 1: Laboratory Screening Experiments

[0032] The data from batch laboratory tests is summarized in Table 1. All polymerizations in Table 1 were carried out in a one liter stainless steel autoclave in 400 cc hexane with 45 cc hexene under 75 psi (517.11 kPa) ethylene head pressure at 60 °C.

[0033] In each case the autoclave was filled with the solvent and hexene and heated under 75 psi (517.11 kPa) of ethylene until equilibrium was reached under rapid stirring. The catalyst was introduced into the reactor under these conditions and the temperature was maintained at 60 °C +/- 5 °C for the duration of the run. The product was recovered and analyzed by GPC and ¹³C NMR spectroscopy to determine the molecular weight and comonomer content. An example of continuous process use of the information contained in Table 1 follows Example 1.1.

Example 1.1

[0034] A one liter mechanically stirred stainless steel autoclave was filled with 400 ml dry and deoxygenated hexane, 45 ml dry and deoxygenated hexane. The reactor was stirred rapidly, pressurized to 75 psi (517.11 kPa) with ethylene, and heated to 60 °C. A catalyst solution in toluene containing 1.42 X 10⁻⁵ moles of r-Me₂Si(Ind)₂HfMe₂ and 4.99 X 10⁻⁵ [PhMe₂NH][B(pfp)₄] was added to the reactor causing a polymerization reaction to occur. After 30 minutes the reactor was vented and the contents were poured into a flask. The solvent was evaporated to yield 5 grams of and

ethylene/hexene copolymer having a $M_w = 151,000$ Daltons, a $M_n = 74,000$ Daltons, and 47.4 wt.% hexene. This represents a productivity of 714 grams PE/gram metallocene.

[0035] The results of the screening experiments collected in Table 1 demonstrate several important trends. These include the following observations.

- Bridged vs. Unbridged Metallocenes
 - + Bridged have 2-3 times the comonomer incorporation (see Cp₂HfMe₂ vs. Et₂SiCp₂HfMe₂ or CpCp*HfMe₂ vs. Me₂SiCpCp*HfMe₂).
 - + Bridged complexes produce higher MW products.
 - + Bridged complexes typically show higher activities.
- [PhMe₂NH][Bpfp)₄] vs. [Ph₃C]Bpfp)₄] vs. [Li•H₂O][Bpfp)₄]
 - + All appear to produce the same polymer at similar rates.
- 15 The Effect of Scavengers on Catalyst Performance
 - + High levels cause lower MW and comonomer content.
 - + Low levels improve productivity without affecting product.
 - + TIBA better than TEAL in terms of effect on product at high levels of scavenger.
 - + high levels depress productivity.
 - Me₂Si(Ind)₂HfMe₂ vrs. Me₂Si(4-Me,2-PhInd)₂HfMe₂
 - + Substituted system shows higher MW even at high comonomer content.
 - + Substituted system shows significant increases in comonomer incorporation.
- 25 The metallocene dichloride route vrs. the metallocene dialkyl route
 - + The chloride free approach produces high MW and comonomer content products.
 - Zr vrs. Hf
 - + Hf systems superior in terms of MW and comonomer incorporation.
 - Mono-Cp Ti vrs. bis-Cp Hf Catalysts
 - + The bis-Cp Hf systems can be modified to have similar performance to the mono-Cp systems.

Part 2: Continuous High Pressure Operation

[0036] The polymerization reactions in Table 2 were performed in a stirred 1.5 L steel autoclave reaction vessel which was equipped to perform continuous Ziegler polymerization reactions at pressures up to 2000 bar and temperatures up to 300 °C. The reaction system was supplied with a thermocouple and pressure transducer to measure temperature and pressure continuously, and with means to supply continuously purified compressed ethylene and 1-butene. Pressure was controlled by pressure reduction valve. Equipment for continuously introducing a measured flow of catalyst solution, and equipment for rapidly venting and quenching the reaction, and of collecting the polymer product were also a part of the reaction system. The ability to add scavenger to the fresh feed prior to the reactor was provided by a high pressure pump. The polymerization was performed with a specified molar ratio of ethylene to comonomer and without the addition of a solvent. The temperature of the reactor containing ethylene and comonomer was equilibrated at the desired reaction temperature. The catalyst solution was prepared by mixing a specified amount of solid metallocene component with the activator component in toluene under an inert atmosphere. This catalyst solution was continuously fed by a high pressure pump into the reactor at a rate which resulted in the desired reactor temperature. The reactor contents were stirred at 1000 rpm and the reactor feed mass flow rate typically used was 40 kg/hr. Exact run conditions including catalyst preparation metallocene component (M) (g), activator component (A) (g), and total volume of solution (L) scavenger aluminum compound to transition metal compound with molar rate (Al:M), catalyst production polymer production rate (kg polymer/hr) average, comonomer molar feed ratio (e.g. C_4/C_2), reactor mass flow rate (kg/hr), and polymer characteristics including melt index (g/10 minutes at 190 °C), and weight percent coinonomer (determined by IR), are also collected in Table 2. All polymerizations were carried out at 225 °C and 1300 bar. The catalyst solutions were prepared using dry and deoxygenated toluene. An example of how to use the information contained in Table 2 follows Example 2.1.

8

5

10

20

30

35

Example 2.1

[0037] Using the reactor design as described above, and using a molar ratio of the ethylene to 1-butene of 0.55 without the addition of a solvent, the temperature of the cleaned reactor containing ethylene and 1-butene was equilibrated at the desired reaction temperature of 225 °C. The catalyst solution was prepared by mixing 0.496 g of solid compound r-Me₂Si(Ind)₂HfMe₂ with 0.9788 activator compound (Ph₃C)+B(pfp)₄ in 20 liter toluene. This catalyst solution was continuously fed by a high pressure pump into the reactor at a rate which resulted in a temperature of 225 °C in the reactor. During this run, ethylene and 1-butene was pressured into the autoclave at a total pressure of 1300 bar. The reactor contents were stirred at 1000 rpm, and the mass feed flow rate through the reactor was held constant at 40 kg/hr. Polymerization was conducted for a 60 to 90 sec. residence time. The yield of polymer product was 6.6 kg/hr. of an ethylene-1-butene copolymer which had a MI of 2.85 and a comonomer incorporation of 13 weight percent butene as measured by IR. Transition metal productivity was calculated at 267 kg polymer/g metallocene.

Table 1

46.8 1944		49.2 1286	2 2			3	3	5	8 8	3 3	3 3	3 2	10 3 10 10 10 10 10 10 10 10 10 10 10 10 10		3 3	3 3 3								
1.8		1.8	1.8	1.8	1.8	1.8	1.8 1.8 2.2 2.1	1.8 1.8 2.2 2.1 1.9	2.1 2.1 2.2 2.2 1.9 1.9	22.2 2.1 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1														2.1 2.1 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1
9 79		8 78					3	12 2	! ! <u>! ! ! ! ! ! !</u>	! ! <u>! ! ! ! ! ! ! </u>	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1		! ! ! ! ! ! ! ! ! ! ! ! ! ! ! ! ! ! ! 	1 1 <u>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 </u>	!	! <u> </u>			/	!	/	/
7 149		9 148												6 6 8 2 8										
30		30				30 30	30 30	30 30 30	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3	30 30 30 30 30 30 30 30 30 30 30 30 30 3
e none	9000	+	+	+	 - - - 	 - - - - 	noue /	<u> </u>	00U 8	non se	3 3	3 3	3 3 5 5	00U E E S		00U 3 3 10U U U U U U U U U U U U U U U U U U U	00	E		00 3 3 100 100	100 000 000 000 000 000 000 000 000 000	00 00 00 00 00 00 00 00 00 00 00 00 00		
7.28E-05 2.50E-06 none	A ROT OR PARTY	4.00L-00	2.91E-05 none	9.10E-05 2.91E-05 none 2.83E-05 1.16E-05 TIBA	2.91E-05 none	2.91E-05 none 1.16E-05 TIBA 3.25E-06 TEAL	2.91E-05 none 1.16E-05 TIBA 3.25E-06 TEAL 7.59E-06 TÉAL	2.91E-05 none 1.16E-05 TIBA 3.25E-06 TEAL 7.59E-06 TEAL 8.68E-06 TEAL	2.83E-05 2.91E-05 inone 2.83E-05 2.91E-05 inone 3.44E-06 3.25E-06 TEAL 7.89E-06 7.59E-06 TEAL 8.90E-06 1.08E-06 TEAL	2.91E-05 none 1.16E-05 TIBA 3.25E-06 TEAL 7.59E-06 TEAL 8.68E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA	2.91E-05 none 1.16E-05 TIBA 3.25E-06 TEAL 7.59E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA	2.91E-05 none 1.16E-05 TBA 3.25E-06 TEAL 7.59E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA	2.91E-05 none 1.16E-05 TIBA 3.25E-06 TEAL 7.59E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA	2.91E-05 none 2.91E-05 none 3.25E-06 TEAL 7.59E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA 7.59E-06 TIBA 7.59E-06 TIBA 7.59E-06 TIBA 7.59E-06 TIBA	2.91E-05 none 2.91E-05 none 3.25E-06 TEAI 7.59E-06 TEAI 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA	2.91E-05 none 2.91E-05 none 2.91E-05 none 3.25E-06 TEAL 3.25E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA	2.91E-05 none 2.91E-05 none 3.25E-06 TEAI 7.59E-06 TEAI 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA	2.916-05 none 2.916-05 none 2.916-05 none 3.256-06 TEAL 3.686-06 TIBA 2.176-06 TIBA 6.516-06 TIBA 7.596-06 TIBA 6.516-06 TIBA 7.596-06 TIBA 8.686-06 TIBA 8.686-06 TIBA	2.91E-05 none 1.16E-05 TIBA 1.25E-06 TEA 1.08E-06 TEA 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA 6.24E-06 TEA 3.58E-06 TEA 3.58E-06 TEA 8.68E-06 TEA 9.99E-06 TEA	3.44E-06 3.25E-06 TEAL 7.89E-06 7.59E-06 TEAL 7.89E-06 7.59E-06 TEAL 7.21E-06 1.08E-06 TIBA 7.22E-06 2.17E-06 TIBA 7.89E-06 7.59E-06 TIBA 7.89E-06 7.59E-06 TIBA 7.89E-06 7.59E-06 TIBA 7.89E-06 3.58E-06 TIBA 7.89E-06 3.58E-06 TEAL 9.34E-06 3.58E-06 TEAL 7.08E-05 8.68E-06 TEAL 7.08E-05 8.68E-06 TEAL 7.08E-05 8.68E-06 TEAL 7.08E-05 8.68E-06 TEAL 7.08E-05 8.59E-06 TEAL	2.91E-05 none 1.16E-05 TIBA 1.25E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA 6.54E-06 TEAL 3.58E-06 TONE 8.68E-06 NONE 8.68E-06 NONE	3.25E-06 TEAL 3.25E-06 TEAL 3.25E-06 TEAL 3.25E-06 TEAL 1.08E-06 TIBA 2.17E-06 TIBA 6.51E-06 TIBA 6.51E-06 TIBA 6.51E-06 TEAL 3.58E-06 TEAL 9.99E-06 TEAL 9.99E-06 TEAL 9.99E-06 OODE 9.99E-06 OODE	3.25E-06 TEAL 3.25E-06 TEAL 3.25E-06 TEAL 3.68E-06 TEAL 3.68E-06 TEAL 4.08E-06 TIBA 5.21E-06 TIBA 6.24E-06 TEAL 3.58E-06 TEAL 3.58E-06 TEAL 3.58E-06 TEAL 3.58E-06 TEAL 3.58E-06 TEAL 3.58E-06 TEAL 4.99E-06 TO ON	9.10E-05 2.91E-05 none 9.10E-05 2.91E-05 none 9.10E-06 3.25E-06 TEAL 7.89E-06 7.59E-06 TEAL 1.21E-06 1.08E-06 TIBA 7.89E-06 7.59E-06 TIBA 6.67E-06 6.51E-06 TIBA 7.89E-06 7.59E-06 None 7.89E-05 8.99E-06 None 7.89E-06 None
7.28E-05 2.		1.46E-US 4.88E-US	9.10E-05 2.91E-05	9.10E-05 4. 2.83E-05 1.	1.46E-05 4. 9.10E-05 2. 2.83E-05 1.	1.46E-US 4.88E-US 9.10E-05 2.91E-05 2.83E-05 1.16E-05 3.44E-06 3.25E-08	1.46E-US 4.88E-US 9.10E-05 2.91E-05 2.83E-05 1.16E-05 3.44E-06 3.25E-06 7.89E-06 7.59E-06	3.44E-06 7.59E-06 7.89E-06 8.90E-05 3.44E-06 7.59E-06 8.90E-06 8.90E-06 8.90E-06 8.90E-06 8.90E-06 8.90E-06 8.90E-06 9.90E-06 9.9	1.40E-US 4. 9.10E-US 2. 2.83E-05 1. 3.44E-06 3. 7.89E-06 7. 8.90E-06 8.	3.44E-06 3.25E-06 3.45E-06 3.25E-06 3.2	3.44E-06 3.25E-06 8.90E-06 1.0E-05 7.89E-06 7.59E-06 8.90E-06 8.68E-06 1.21E-06 1.08E-06 6.72E-06 6.11E-06	3.44E-06 3.25E-06 3.2	3.44E-06 3.25E-06 3.24E-06 3.25E-06 3.2	3.44E-06 3.25E-06 3.25E-06 3.25E-06 3.25E-06 3.35E-06 3.3	3.44E-06 3.1.48E-06 3.	1.40E-US 4.88E-US 9.10E-US 2.91E-US 2.83E-US 1.16E-US 3.44E-06 3.25E-US 1.22E-US 1.08E-US 1.22E-US 2.22E-US 2.22E-US 2.22E-US 2.34E-US 3.58E-US 3.58E-US 3.58E-US 2.65E-US 8.68E-US 3.58E-US 3.5	1.40E-US 4. 9.10E-US 2. 2.83E-US 1. 1.21E-US 1. 1.21E-US 1. 1.21E-US 1. 1.87E-US 2. 1.87E-US 3. 1.87E-US 3. 1.87E-US 3. 1.87E-US 3. 1.87E-US 3. 1.87E-US 3. 1.87E-US 3.	1.40E-05 4. 9.10E-05 2. 2.83E-05 1. 3.44E-06 3. 8.90E-06 8. 8.90E-06 8. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 3. 9.34E-05 8. 2.65E-05 8.	1.40E-05 4. 9.10E-05 2. 2.83E-05 1. 3.44E-06 3. 7.89E-06 7. 1.21E-06 1. 1.87E-05 6. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 8. 7.89E-06 7. 7.89E-06 8. 7.89E-06 8. 7.89E	1.40E-05 4. 9.10E-05 2. 2.83E-05 1. 3.44E-06 3. 3.44E-06 3. 1.21E-06 1. 1.87E-05 6. 1.87E-05 6. 1.87E-05 6. 1.87E-05 6. 1.87E-05 6. 2.65E-05 8. 3.20E-05 3. 3.20E-05 3.	1.40E-05 4. 2.83E-05 1. 2.83E-06 3. 3.44E-06 3. 2.22E-06 2. 1.87E-06 6. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 8. 7.89E-06 9. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 8. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 8. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 7. 7.89E-06 8. 7.89E-06 7. 7.89E-06 8. 7.89E-06 8. 7.89E	1.40E-US 4.88E-US 9.10E-05 2.91E-05 2.83E-06 1.16E-05 7.89E-06 7.59E-06 8.90E-06 8.68E-06 1.21E-06 1.08E-06 6.67E-06 2.17E-06 6.67E-06 6.51E-06 7.89E-06 7.59E-06 9.34E-06 3.58E-06 9.34E-05 9.99E-06 9.35E-05 8.68E-06 9.35E-05 8.68E-06	3.44E-06 3.25E-06 5.29E-06 5.29E-06 5.29E-06 5.29E-06 5.29E-06 5.29E-06 5.22E-06 5.2	2.65E-05 3. 2.0E-05 3. 1.44E-05 3. 1.44E-05 3. 2.0E-05 3. 1.44E-05 3. 2.0E-05 3. 1.44E-05
J.	[Ph,C][B(p(p),]												Tr@j	P(d)c	Prdja	Pr@ja	Pr@jc	Pr@g	[Fid]c	[7]	[76]c	[7d]c	LIB(pfp)4 • H ₂ O LIB(pfp)4 • H ₂ O LIB(pfp)4 • H ₂ O [Ph ₂ C][B(pfp)4 [Ph ₂ C][LIB(pfp), +h2O LIB(pfp), +h2O IPh2CIB(pfp), Ph3CIB(pfp), Ph3CIB(pfp),
	1.2 r-Me,Si(Ind),HfMe,																nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HICI,	nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe, nd),HIMe,	nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHICIz	nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHICIz nd)zHICIz nd)zHICIz	nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHICIz nd)zHICIz nd)zHICIz nd)zHICIz	nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHICIz nd)zHICIz nd)zHICIz nd)zHICIz nd)zHICIz	nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz nd)zHICiz	nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez nd)zHIMez
1.1	1.2 6		1.3	1.3	1.3	1.3 1.4 Comp 1.5	1.3 r Comp 1.5 r Comp 1.6 r	Comp 1.5 Comp 1.5 Comp 1.7 I	Comp 1.5 Comp 1.5 Comp 1.8 Com	Comp 1.5 Comp 1.6 Comp 1.9 Com	Comp 1.5 Comp 1.6 Comp 1.7 Comp 1.8 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.10 Comp	Comp 1.5 Comp 1.6 Comp 1.8 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.10 Com	Comp 1.5 Comp 1.6 Comp 1.8 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.12 Com	Comp 1.3 Comp 1.4 Comp 1.9 Comp 1.9 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.13	Comp 1.3 Comp 1.4 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.12 Comp 1.13 C	Comp 1.5 Comp 1.6 Comp 1.9 Comp 1.9 Comp 1.9 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.11 Comp 1.13 Comp 1.14 C	Comp 1.5 Comp 1.10 Comp 1.11 Comp 1.	Comp 1.5 rMe ₂ Si(I) Comp 1.5 rMe ₂ Si(I) Comp 1.7 rMe ₂ Si(I) Comp 1.9 rMe ₂ Si(I) Comp 1.9 rMe ₂ Si(I) Comp 1.10 rMe ₂ Si(I) Comp 1.10 rMe ₂ Si(I) Comp 1.11 rMe ₂ Si(I) Comp 1.12 rMe ₂ Si(I) Comp 1.13 rMe ₂ Si(I) Comp 1.14 cp ₂ HIMe	Comp 1.5 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.11 Comp 1.13 Comp 1.	Comp 1.5 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.11 Comp 1.13 Comp 1.14 Comp 1.15 Comp 1.	Comp 1.5 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.12 Comp 1.13 Comp 1.14 Comp 1.15 Comp 1.15 Comp 1.14 Comp 1.15 Comp 1.14 Comp 1.15 Comp 1.14 Comp 1.14 Comp 1.15 Comp 1.14 Comp 1.15 Comp 1.14 Comp 1.	Comp 1.5 Comp 1.6 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.13 Comp 1.13 Comp 1.13 Comp 1.13 Comp 1.13 Comp 1.15 Comp 1.1	Comp 1.5 Comp 1.5 Comp 1.5 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.12 Comp 1.13 Comp 1.15	Comp 1.5 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.10 Comp 1.11 Comp 1.11 Comp 1.12 Comp 1.13 Comp 1.15 Comp 1.

* In-situ activation, dual functioning of alkylaluminum. Comp'' indicates comparative example.

Exp		Weight	Activator (A)	Welght	Weight Volume	M:A	Scav	A1:M	. ;		Productio	%¥ ¥	wt% KaPE/
3 2	Metallocene (M)	M (gr)		A (gr)	Toluen	Mole			ϋ	₹	n Rate	ŭ	Mo
					e (L)	Ratio					(Kg/hr)		,
2.1	2.1 r-Me ₂ Si(Ind) ₂ H/Me ₂	0.496	trityl B(pfp),-	0.978	20	0.94	TOA	33	33 0.55	2.85	9.9	13	287K
2.2	2.2 r-Me ₂ Si(Ind) ₂ HfMe ₂	0.552	0.552 DMAH'B(pfp),	0.957	20	0.93	TOA	34	0.55	3.02	6.1	13.7	251K
2.3	2.3 r-Me ₂ Si(Ind) ₂ HfMe ₂	0.662	DMAH'B(pfp)4"	0.932	20	+	TOA	32		25.1	6.9	25.4	178X
2.4	٤	0.891	0.891 DMAH*B(pfp),*	1.063	20	0.99	TOA	55	-	63.3	ď	A 38 5	200K
	Me, Si(2Me, 4Phlnd), HIMe,							3	•	?	•	2	70c7
2.5 6-	٤	0.891	DMAH'B(pfp)4"	1.063	20	-	TO A	6	0.6	9.95	8.9	28 S	SSK
	Me ₂ Si(2Me,4PhInd) ₂ HfMe ₂										?	?	<u></u>
2.6	2.6 r-Me,Si(2MInd),H(Me,	0.497	0.497 DMAH'B(p(p),-	0 86	10	-	40,		97.0	100		1	

Part 3: Continuous High Temperature Solution Process

30

35

40

45

50

55

[0038] The polymerization reaction was performed in a stirred, liquid filled 2 L jacketed steel reactor equipped to perform continuous insertion polymerization in presence of an inert C_6 hydrocarbon (naphta) solvent at pressures up to 120 bar and temperatures up to 240 °C. The reactor was typically stirred at 1000 rpm during the polymerization. The reaction system was supplied with a thermocouple and a pressure transducer to monitor changes in temperature and pressure continuously, and with means to supply continuously purified ethylene, 1-octene, and solvent. In this system, ethylene dissolved in the hydrocarbon solvent, 1-octene, tri-n-octyl aluminum (TOA) used as a scavenger, and optionally H2, are pumped separately, mixed, and fed to the reactor as a single stream, refrigerated to -40 °C using liquid NH $_3$ as a coolant. The transition metal component (TMC) was dissolved in a solvent/toluene mixture (9/1 vol/vol) whereas the non-coordinating anion (NCA) activator was dissolved in toluene or slurried in the hydrocarbon solvent. Both components were pumped separately, mixed at ambient temperature, and cooled to -1 °C prior to entering the reactor. The reactor temperature was set by adjusting the temperature of an oil bath used as a reservoir for the oil flowing through the reactor wall jacket. Next, the polymer molecular weight (MW) or MI was controlled independently by adjusting the ethylene conversion (% C_2) in the reactor via the catalyst flow rate. Finally, the polymer density was controlled by adjusting the ethylene/1-octene weight ratio in the feed.

[0039] The results of ten polymerization experiments performed in the reactor described above with three different TMC's (A = rac-dimethylsilyl bis(1-indenyl) hafnium dimethyl, B = dimethylsilyl bis(2-methyl-4-phenyl-1-indenyl) hafnium dimethyl, and C = dimethylsilyl (tetramethylcyclopentadienyl)butylamino titatium dimethyl) and [PhMe₂NH]B(pfp)₄] used as the activator (NCA) are summarized in Table 1. For example, the polymerization reaction in run #1 was carried out at 150 °C and 86.4 bar with A. Solvent, ethylene, and 1-octene were continuously fed to the reactor a; a rate of 7 kg/hr., and 0.53 kg/hr., respectively, The 1-octene/ethylene weight ratio in the feed was 0.431 in this case. 3.5 mg/hr. of A dissolved in a 9/1 solvent/toluene (vol/vol) mixture and 4.9 mg/hr. of NCA dissolved in toluene were continuously fed to the reactor, resulting in an ethylene conversion of 80.5% and a 1-octene conversion of 43%. For a reactor residence time of 8 minutes, the polymer yield was 1.22 kg/hr, under these conditions. By adding TOA at a rate of 0.08 mmol/hr. to the feed, the catalyst productivity lined out around 348 kgPE/g A. This experiment resulted in an ethylene/1-octene copolymer containing 18.77 weight % comonomer (FTIR) with a weight-average MW of S8.000 g/mol (GPC), a polydispersity index of 2.1 (GPC), a MI of 1.4 dg/min., a density of 0.8991 g/cc, and a MIR (121.6/12) of 34.

[0040] At constant reactor temperature and feed composition, the polymer MW can be adjusted in this low pressure/ high temperature polymerization process by controlling the ethylene conversion in the reactor via the catalyst flow rate; typically, the higher the ethylene conversion, the lower the polymer MW (the higher the MI). The above examples showed that surprisingly both bridged bis (Cp) hafnocenes (A and B) exhibit superior MW and comonomer incorporation capability than the bridged mono(Cp) titanocene (C) does. For example, at 140 °C and with a 1-octene/ethylene weight ratio in the feed equal to about 0.43 to 0.44, A and B produced, respectively, a 1.6 MI/0.8937 g/cc copolymer at 87.9% ethylene conversion (run #3) and a 0.62 MI/0.8880 g/cc copolymer at 80.7% ethylene conversion (run #6), whereas (C) produced a 1.5 MI/0.9035 g/cc copolymer at a substantially lower ethylene conversion of 77.1% (run #9). Similarly, at 130 °C and with a 1-octene/ethylene weight ratio in the feed equal to about 0.93 to 0.94, A produced a 1.23 MI/0.8761 g/cc copolymer at 73.9% ethylene conversion (run #4), whereas (C) made a 3.2 MI/0.8866 g/cc copolymer at a lower ethylene conversion of 69.2% (run #10).

Table 3

	-	_	_	_	_	_	-,-		.		_
CO/CO	ka/hr	0.431	7670	27.0	0.000	2450	0.421	0.420	0.430	0.425	7,000
౮	ka/hr	0.530	0 530	0 530	1 440	2 2 2	0.3.0	0.320	0.320	0 520	1 140
ర	kg/hr	123	123	2 2	12/2	125	1 2 5	1 2 2	127	122	123
اد اگ 30	kg/hr	7.0	6.9	7.1	8.0	7.0	2.5	3 -	7.0	7.0	6.9
۵	bar	86.4	86.3	86.6	86.9	86.4	AR A	86.5	86.5	86.7	86.6
±	°C	150.0	159.9	142.6	130.3	149.1	139.9	159.2	150.5	140.9	131.5
AIR3	mmol/hr	0.08	0.08	0.11	0.11	0.08	0.08	0.08	0.08	0.08	0.10
ACT/TM	тој/то	0.86	0.88	0.82	0.88	1.03	1.03	1.02	0.99	1.00	1.01
ACT	mg/hr	4.9	3.7	5.3	2.0	6.7	6.1	7.2	6.0	5.1	4.4
Ϋ́	mg/hr	3.5	2.6	4.0	1.4	5.5	5.0	2.9	2.5	2.1	1.8
MW (TM)	g/mol	494.5	494.5	494.5	494.5	674.5	674.5	329.9	329.9	329.9	329.9
Ž.		4	٧	٧	٧	89	8	ပ	ပ	U	U
# 4		-	7	9	4	2	9	7	80	5	Ē

_					_	_	_		_	_	_		_	_	_		_
100		MWINI		2.1		2.1	,	3		9 6		3.3	2.5		2.5	1	7.7
Mer	lom/o	IOIII/A		88000 00	00000	000000	8400000	00000		95000 00			91000.00	20000	91000.00	970000	2000
Ceinc	% JW			18.77	46.30	10.30	21 49	30.69	30.00	23.65	26.05	20.03	12.22	42.00	13.30	15.66	25.03
density	a/cc			0.8991	0 9038	2000	0.8947	0.8761	10.00	0.8915	OBBBO		0.9121	0 9082	0.3002	0.9035	O Bee
MIR	121.6/12			34.0	32.2		53.7	316		20.0	65.8		23.8	25.3		26.7	24.5
Ψ	dg/min			1.40	1.20		1.60	1.23		1.40	0.62		2.10	1.90		1.50	3.20
cat prod	kg PE/g TM			348	413		336	922		212	266	25.0	320	422		531	639
yield	kg/hr			1.22	1:047		1.35	1.29		1:1	1.33	900	S	1.05		71.1	1.15
ž	min			8.0	8.0	į	F. /	7.9		9.1	7.8	7.0		8.0 0.8		9.1	7.8
రీ	% conv			43.0	33.1		94.9	34.8	3	93.9	66.5	22.5	7	28.3		-33.	26.3
°	% conv		1	80.5	73.1	1	2/2	73.9		2.0	2 08	69.0		738		-!	69.2

(1) For all runs the catalyst activator was [DMAH][B(pfp)4] and the scavenger was tri-n-octyl aluminum. 5 10 15 20 25 30 Other abbreviations above include: C₆ = hydrocarbon (naphta)solvent $B = Me_2Si(2-Me-4Phlnd)_2IIfMe_2$ C = Me₂SiCp*NbutylTiMe₂ A = r-Me₂Si(Ind)₂HfMe₂ 35 $C_{\mathbf{l}} = 1$ -octene C_2 = ethylene 40 45

Claims

- 50 1. A method for insertion polymerization of olefinically or acetylenically unsaturated monomers comprising contacting one or more of said monomers under suitable polymerization process conditions with an ionic catalyst composition comprising the reaction product of a catalytically suitable Group 4, 5, 6 or 8 transition metal compound having a univalent hydride, alkyl or silyl ligand and a hydrated salt comprising a Group 1 or 2 cation and a non-coordinating anion.
 - 2. A method according to Claim 1 in which the hydrated salt comprises a lithium cation.
 - 3. A method according to Claim 2 in which non-volatile LiOH is produced and removed from any monomer and any

diluent prior to recycle.

- 4. A method according to any of the preceding claims in which the transition metal compound is a bridged, biscy-clopentadienyl hafnium compound.
- 5. Use as an anion precursor for reacting with a catalytically suitable Group 4, 5, 6 or 8 transition metal compound having a univalent hydride, alkyl or silyl ligand, of a hydrated salt comprising a Group 1 or 2 cation and a non-coordinating anion.



Europäisches Patentamt European Patent Office Office européen des brevets



(11) EP 1 063 244 A3

(12)

EUROPEAN PATENT APPLICATION

(88) Date of publication A3: 26.11.2003 Bulletin 2003/48

(51) Int CI.7: **C08F 210/18**, C08F 2/06, C08F 4/645

(43) Date of publication A2: 27.12.2000 Bulletin 2000/52

(21) Application number: 00121154.9

(22) Date of filing: 18.12.1996

(84) Designated Contracting States: BE DE ES FR GB IT NL

(30) Priority: 19.12.1995 US 8893 P

(62) Document number(s) of the earlier application(s) in accordance with Art. 76 EPC: 96944998.2 / 0 868 445

(71) Applicant: ExxonMobil Chemical Patents Inc. Baytown, TX 77520-5200 (US)

(72) Inventors:

Turner, Howard W.
 Campbell, CA 95008 (US)

Speed, Charles S.
 Dayton, Texas 77535 (US)

Folle, Bernard J.
 1380 Lasne (BE)

Crowther, Donna J.
 Seabrook, Texas 77586 (US)

Walzer, John F.
 Seabrook, Texas 77586 (US)

Fisher, Richard A.
 Malvern, Pennsylvania 19355 (US)

 Vaughan, George A. Houston, Texas 77025 (US)

(74) Representative:

Veldhulzen, Albert Dirk Willem et al Exxon Chemical Europe Inc., P.O.Box 105 1830 Machelen (BE)

(54) High temperature olefin polymerization process

(57) The invention provides a method for insertion polymerization of olefinically or acetylenically unsaturated monomers comprising contacting one or more of said monomers under suitable polymerization process conditions with an ionic catalyst composition comprising the

reaction product of a catalytically suitable Group 4, 5, 6 or 8 transition metal compound having a univalent hydride, alkyl or dilyl ligand and a hydrated salt comprising a Group 1 or 2 cation and a non-coordinating anion.



EUROPEAN SEARCH REPORT

Application Number EP 00 12 1154

	DOCUMENTS CONSIDE	RED TO BE RELEVANT	,	
Category	Citation of document with inc of relevant passag		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
P,X	PROCESS" RESEARCH DISCLOSURE PUBLICATIONS, HAMPSI	HIRE, GB, 1996 (1996-08-01), pages 1	1-5	C08F210/18 C08F2/06 C08F4/645
x	EP 0 658 576 A (TOS 21 June 1995 (1995- * page 2, line 55 - * page 3, line 54 * * page 5, line 38 - * page 7; example 4	96-21) page 3, line 30 * line 39 *	1,2,4,5	
X	US 5 408 017 A (FOL 18 April 1995 (1995 * column 13, line 2		1-5	
A	EP 0 582 268 A (TOS 9 February 1994 (1992) * page 3, line 2 - * page 3, line 15 - * page 6; example 4	94-02-09) line 3 * line 17 *	1-5	TECHNICAL FIELDS SEARCHED (Int.Ct.7)
A	EP 0 598 134 A (IDE) 25 May 1994 (1994-0) * example 1 *		1-5	<u>.</u>
	The present search report has b	een drawn up for all claims Date of completion of the search		Examiner
	MUNICH	16 September 2003	3 Tho	mas, D
X : parti Y : parti docu A : tech O : non	ATEGORY OF CITED DOCUMENTS culturly relevant if bornbined with anothment of the same category nological background written disclosure mediate document	T : theory or principle E : earlier patent door after the filing date er D : document ofted in L : doournent cited to	underlying the ir ument, but publis the application r other reasons	ivention thed on, or

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 00 12 1154

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

16-09-2003

Patent document cited in search report	Publication date		Patent family member(s)	Publication date
EP 0658576 A	21-06-1995		59412047 D1 59412047 T2 0658576 A1 3033452 B2 7224106 A 5830820 A	03-09-1998 15-04-1999 21-06-1995 17-04-2000 22-08-1995 03-11-1998
US 5408017 A	18-04-1995	US US CA DE DE EP ES JP WOS US US US US US AR AR AR AR AR DE DE DE DE DE DE DE DE DE DE DE DE DE	5278119 A 5198401 A 2117888 C 69322865 D1 69322865 T2 0646140 A1 2126653 T3 3058690 B2 7508545 T 9325590 A1 5407884 A 5621126 A RE37788 E1 5483014 A 6423795 B1 147761 T 9106589 A 2085581 A1 19122275 D1 69122275 T2 59124255 D1 69124255 T2 551277 T3 0551277 T3 0670334 A2 0672689 A1 0670334 A2 0672689 A1 0672688 A1 2079330 T1 2092913 T3 2954351 B2 5507756 T 123370 B1 2118203 C1 9200333 A2 66294625 B1 6355592 B1 5801113 A	11-01-1994 30-03-1993 15-05-2001 11-02-1999 17-06-1999 05-04-1995 01-04-1995 01-04-1995 23-12-1993 18-04-1997 09-07-2002 09-01-1996 23-07-2002 15-02-1997 01-06-1993 23-12-1991 24-10-1996 15-05-1997 27-02-1997 05-06-1997 10-10-1996 14-07-1997 21-07-1993 06-09-1995 20-09-1995 20-09-1995 20-09-1995 20-09-1995 20-09-1995 21-07-1997 21-07-1993 06-09-1995 20-09-1999

For more details about this annex: see Official Journal of the European Patent Office, No. 12/82

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 00 12 1154

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

16-09-2003

AU 1294588 A 24-08-1986 BR 8805025 A 17-10-1986 CA 1338687 C 29-10-1996 CA 1340136 C 17-11-1996 DE 3855727 D1 06-02-1997 OK 548988 A 30-11-1986 EP 0277003 A1 03-08-1986 EP 0478913 A1 08-04-1995 ES 2095273 T3 16-02-1997 FI 884486 A 29-09-1986 EP 0582268 A 09-02-1994 JP 6049122 A 22-02-1997 CA 2101805 A1 05-02-1997 DE 69322139 D1 24-12-1998 DE 69322139 T2 06-05-1997 DE 69322139 T2 06-05-1997 US 5407882 A 18-04-1999 JP 2988226 B2 13-12-1998 JP 2988226 B2 13-12-1999 JP 2988226 B2 13-12-1999 JP 6228222 A 16-08-1997 EP 0598134 A 25-05-1994 JP 5331228 A 14-12-1999 JP 3299275 B2 08-07-2096 CA 2113806 A1 09-12-1993	AU 1294588 A 24-08-1988 BR 8805025 A 17-10-1989 CA 1338687 C 29-10-1996 CA 1340136 C 17-11-1998 DE 3855727 D1 06-02-1997 DK 548988 A 30-11-1988 EP 0277003 A1 03-08-1988 EP 0478913 A1 08-04-1992 ES 2095273 T3 16-02-1997 FI 884486 A 29-09-1988 EP 0582268 A 09-02-1994 JP 6049122 A 22-02-1994 CA 2101805 A1 05-02-1994 CA 2101805 A1 05-02-1994 DE 69322139 D1 24-12-1998 DE 69322139 T2 06-05-1999 DE 69322139 T2 06-05-1999 US 5407882 A 18-04-1995 JP 2988226 B2 13-12-1999 JP 6228222 A 16-08-1994 EP 0598134 A 25-05-1994 JP 5331228 A 14-12-1993 EP 0598134 A1 25-05-1994 JP 3299275 B2 08-07-2002	Patent document cited in search report	t	Publication date		Patent fami member(s		Publication date
JP 6056927 A 01-03-1994 CA 2101805 A1 05-02-1994 DE 69322139 D1 24-12-1998 DE 69322139 T2 06-05-1999 EP 0582268 A2 09-02-1999 US 5407882 A 18-04-1999 JP 2988226 B2 13-12-1999 JP 6228222 A 16-08-1994 EP 0598134 A 25-05-1994 JP 5331228 A 14-12-1993 EP 0598134 A1 25-05-1994 JP 3299275 B2 08-07-2002 CA 2113806 A1 09-12-1993	JP 6056927 A 01-03-1994 CA 2101805 A1 05-02-1994 DE 69322139 D1 24-12-1998 DE 69322139 T2 06-05-1999 EP 0582268 A2 09-02-1994 US 5407882 A 18-04-1995 JP 2988226 B2 13-12-1999 JP 6228222 A 16-08-1994 EP 0598134 A 25-05-1994 JP 5331228 A 14-12-1993 EP 0598134 A 25-05-1994 JP 331228 A 14-12-1993 EP 0598134 A1 25-05-1994 JP 3299275 B2 08-07-2002 CA 2113806 A1 09-12-1993 W0 9404578 A1 03-03-1994	US 5408017	A		AU BR CA CA DE DK EP EP ES	1294588 8805025 1338687 1340136 3855727 548988 0277003 0478913 2095273	A A C C D1 A A1 A1 T3	24-08-1988 17-10-1989 29-10-1996 17-11-1998 06-02-1997 30-11-1988 03-08-1988 08-04-1992 16-02-1997
JP 48054243 A 30-07-1973 EP 0598134 A1 25-05-1994 JP 3299275 B2 08-07-2002 CA 2113806 A1 09-12-1993	JP. 48054243 A 30-07-1973 EP 0598134 A1 25-05-1994 JP 3299275 B2 08-07-2002 CA 2113806 A1 09-12-1993 W0 9404578 A1 03-03-1994	EP 0582268	A 09	9-02-1994	JP CA DE DE EP US JP	6056927 2101805 69322139 69322139 0582268 5407882 2988226	A A1 D1 T2 A2 A B2	01-03-1994 05-02-1994 24-12-1998 06-05-1999 09-02-1994 18-04-1995 13-12-1999
	··	EP 0598134		5-05-1994	JP. EP JP CA WO	48054243 0598134 3299275 2113806 9404578	A1 B2 A1 A1	30-07-1973 25-05-1994 08-07-2002 09-12-1993 03-03-1994

b For more details about this annex: see Official Journal of the European Patent Office, No. 12/82

This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

□ BLACK BORDERS
□ IMAGE CUT OFF AT TOP, BOTTOM OR SIDES
□ FADED TEXT OR DRAWING
□ BLURRED OR ILLEGIBLE TEXT OR DRAWING
□ SKEWED/SLANTED IMAGES
□ COLOR OR BLACK AND WHITE PHOTOGRAPHS
□ GRAY SCALE DOCUMENTS
□ LINES OR MARKS ON ORIGINAL DOCUMENT
□ REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY

IMAGES ARE BEST AVAILABLE COPY.

OTHER:

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.